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# **ORIGINAL ARTICLE** The *MLL* recombinome of acute leukemias in 2017

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Chromosomal rearrangements of the human *MLL/KMT2A* gene are associated with infant, pediatric, adult and therapy-induced acute leukemias. Here we present the data obtained from 2345 acute leukemia patients. Genomic breakpoints within the *MLL* gene and the involved translocation partner genes (TPGs) were determined and 11 novel TPGs were identified. Thus, a total of 135 different *MLL* rearrangements have been identified so far, of which 94 TPGs are now characterized at the molecular level. In all, 35 out of these 94 TPGs occur recurrently, but only 9 specific gene fusions account for more than 90% of all illegitimate recombinations of the *MLL* gene. We observed an age-dependent breakpoint shift with breakpoints localizing within *MLL* intron 11 associated with acute lymphoblastic leukemia and younger patients, while breakpoints in *MLL* intron 9 predominate in AML or older patients. The molecular characterization of *MLL* gene with clinical outcome. This study provides a comprehensive analysis of the *MLL* recombinome in acute leukemia and demonstrates that the establishment of patient-specific chromosomal fusion sites allows the design of specific PCR primers for minimal residual disease analyses for all patients.

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# INTRODUCTION

Chromosomal rearrangements involving the human MLL gene are recurrently associated with the disease phenotype of acute leukemias.<sup>1,2</sup> The presence of distinct *MLL* rearrangements is an independent dismal prognostic factor, while very few MLL rearrangements display either a good or an intermediate outcome.<sup>3,4</sup> It became also clear from recent studies that the follow-up of patients during treatment and therapy adjustment based on minimal residual disease (MRD) monitoring has a very strong impact on outcome.<sup>5-7</sup> For this purpose, we established a diagnostic network that allowed different study groups and clinical centers to obtain genomic MLL breakpoint sequences that can be directly used for quantifying MRD levels in patients. The current work flow to identify MLL rearrangements includes a prescreening step (cytogenetic analyses,<sup>8,9</sup> split-signal fluores-cence *in situ* hybidization<sup>10–12</sup> or reverse-transcription PCR) in combination with long-distance inverse PCR that was performed on small amounts (~1 µg) of isolated genomic DNA.<sup>13</sup> This allowed us to readily identify reciprocal translocations, complex chromosomal rearrangements, gene internal duplications, deletions or inversions on chromosome 11g, and MLL gene insertions into other chromosomes, or vice versa, the insertion of partner chromosome material into the MLL gene located at 11q23.

To gain insight into the frequency of distinct *MLL* rearrangements, all prescreened samples of infant, pediatric and adult leukemia patients were sent for analysis to the Frankfurt Diagnostic Center of Acute Leukemia (DCAL) after cytogenetic prescreening. All prescreened *MLL* rearrangements were successfully analyzed at the Frankfurt DCAL, and patient-specific *MLL* fusion sequences for MRD monitoring were obtained.

## MATERIALS AND METHODS

#### Patient material

Genomic DNA was isolated from bone marrow and/or peripheral blood samples of leukemia patients and sent to the DCAL (Frankfurt/Main, Germany). Patient samples were obtained from different study groups (the AMLCG-study group, Munich; the GMALL study group, Berlin; Polish Pediatric Leukemia and Lymphoma Study Group; Zabrze; I-BFM network) and diagnostic centers in Europe (Aarhus, Berlin, Barcelona, Bordeaux, Bratislava, Brest, Bristol, Catania, Copenhagen, Ekaterinburg, Frankfurt, Giessen, Granada, Graz, Grenoble, Haifa, Hamburg, Hanover, Heidelberg, Jena, Jerusalem, Kiel, Lille, Lisbon, Madrid, Minsk, Montpellier, Monza, Moscow, Munster, Munich, Nancy, Nantes, Newcastle upon Tyne, Olomouc, Padua, Paris, Porto, Prague, Reims, Rotterdam, Strasbourg, Tampere, Tel Hashomer, Toulouse, Turku, Ulm, Valenciennes, Vienna, Zabrze and Zurich) or centers located outside of Europe (Adelaide, Boston, Brisbane, Buenos Aires, Hong Kong, Houston, Rio de Janeiro, Seoul, Sydney and Tohoku), where acute leukemia patients are enrolled in local study groups. Informed consent was obtained from all patients or patients' parents/legal guardians, and control individuals.

#### Long distance inverse PCR experiments

All DNA samples were treated and analyzed as described.<sup>13–16</sup> Briefly, 1  $\mu$ g genomic patient DNA was digested with restriction enzymes and re-ligated to form DNA circles before long-distance inverse PCR analyses. Restriction polymorphic PCR amplimers were isolated from the gel and subjected to DNA sequence analyses to obtain the patient-specific fusion sequences. This genomic DNA fusion sequence is idiosyncratic for each leukemia patient and was made available to the sender of the DNA sample. The average processing time was around 5 working days.

## Data evaluation and statistical analyses

All clinical and experimental patient data were implemented into a database program (FileMaker Pro, FileMaker Inc., Santa Clara, CA, USA) for further analysis. Information about all individual patients was used to compare all defined subgroups and to perform statistical analyses to retrieve important information or significant correlations.  $\chi^2$ -tests were performed to identify significant deviations from mean values.

#### Nomenclature

We are well aware about all the changes in the HUGO gene nomenclature over the past years. However, for the readability of the text, we use the following gene nomenclature throughout the text: *MLL* (*KMT2A*); *AF4* (*AFF1*); *LAF4* (*AFF3*); *AF5* (*AFF4*); *ENL* (*MLLT1*); *AF9* (*MLLT3*); *AF6* (*MLLT4*); *AF17* (*MLLT6*); *AF10* (*MLLT10*); and *AF1Q* (*MLLT11*).

# RESULTS

#### The study cohort

To analyze the recombinione of the human MLL gene, 2381 prescreened acute leukemia samples were obtained from the above-mentioned centers from 2003 to 2016. In all cases, we first used PCR experiments combined with sequence analysis to diagnose the direct MLL fusion allele, and in case of failure or having a complex rearrangement, the reciprocal MLL fusion allele was analyzed. Successful analysis could be performed for all patient samples. In 31 cases we were only able to characterize the reciprocal MLL fusion allele to guarantee subsequent MRD experiments. Complete data were available on 2345/2381 cases (gender, age at diagnosis, disease type and subtype, or information about de novo or secondary leukemia). Genetic and clinical information of these 2345 patients are summarized in Table 1. The 36 excluded cases had the following *MLL* rearrangements:  $9 \times$ MLL-AF9; 8 × MLL-AF4; 4 × MLL-ENL; 4 × MLL-AF10; 3 × MLL-AF6; 2× MLL-AF17; 2× MLL-EPS15; 1× MLL-GAS7; 1× MLL-LOC100128568; 1 × MLL–CREBBP; and 1 × MLL–PTD. The exclusion of these 36 patients did not affected with the general conclusions made in this study.

## Age distribution according to clinical subtypes

We first analyzed our cohort according to the age at diagnosis. As displayed in Figure 1, the age distribution is quite similar to the expected distributions known from other cancer registries. MLL-r acute lymphoblastic leukemia (ALL) incidence has a peak in the first 2 years, then declines during the pediatric and young adult phase and then steadily increases again with age. A similar picture was observed with MLL-r acute myeloid leukemia (AML) patients, however, missing the postnatal peak seen for infant ALL. For the purpose of our study we separated our cohort into an 'infant acute leukemia cohort' (0.03–12 months; n = 876: 692 ALL, 160 AML and 24NA) (not annotated), a 'pediatric acute leukemia group' (>12 months-18 years; n=671: 313 ALL, 339 AML and 19NA) and an 'adult acute leukemia patient' group (>18 years; n = 798: 415 ALL, 373 AML and 10NA). As shown in Figure 1, we also added information about therapy-induced leukemia (n = 110). Fifty-three patients could not be categorized into 'ALL' or 'AML' because they received other diagnoses (mixed lineage leukemia (MLL) = 38, myelodysplastic syndrome = 7 and lymphoma = 4), or because we had no information from the corresponding center (unknown disease type = 4).

Identification of MLL rearrangements and their distribution in clinical subgroups

The most frequent *MLL* rearrangements in these six subgroups were summarized in Figure 2. Infant ALL (n = 692) displayed 338 *AF4*, 113 *AF9*, 154 *ENL*, 39 *AF10*, 1 *AF6* and 16 *EPS15* gene fusions. Additional 31 *MLL* rearrangements were identified (*9p13.3*, *11q23.3*, *ACER*, *AF1Q*, *LAF4*, *AF5*, *BTBD18*, *CLTA*, *DCP1A*, *EEFSEC*, 14 cases (NA) with no der(11) and only a reciprocal fusion allele, *PICALM*, *PRPF19* and *TNRC18*).

Infant AML (*n* = 160) was represented by 4 *AF4*, 40 *AF9*, 2 *ENL*, 43 *AF10*, 24 *ELL*, 2 *AF6* and 1 *EPS15* gene fusion. Another 44 *MLL* rearrangements have been characterized (*11q24*, *ABI1*, *ABI2*, *AF1Q*, *FLNA*, *FNBP1*, *GAS7*, *KIAS1524*, *MYO1F*, 14 cases (NA) with no der (11) and only a reciprocal fusion allele, *NEBL*, *NRIP3*, *PICALM*, *SEPT5*, *SEPT6* and *SEPT9*).

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Table 1.	Overview about all i	nvestigated	TPGs								
#	Direct TPG		Infant			Pediatric			Adult		Total
		ALL	AML	Other	ALL	AML	Other	ALL	AML	Other	
1	AFF1/AF4	338	4	10	139	3	10	332	3	_	839
2	MLLT3/AF9	113	40	5	56	132	3	9	90	1	449
3	MLLT1/ENL	154	2	4	56	21	1	50	14		302
4	MLLT10/AF10	39	43	2	12	66	1	1	33		197
5	PTD	—			—	6	—	1	98	2	107
6	ELL	_	24	1		24	_	1	45	2	97
	MLL14/AF6	1	2	1	16	28		9	38	1	95
8	EPS15	10	12	I	6	5	_	4	5	_	38
10	MILLITI/AFTQ	14	13	1	2	7	_	_	2		23
10	SEPT6	14	5		5	10	_	_	2		17
12	MUT6/AF17	_		_	1	2	_	_	11		14
13	SEPT9	_	2	_	_	5	_	_	6	_	13
14	AFF3/LAF4	3	_	_	5	_	_	_	_	_	8
15	TET1					1	_	2	3		6
16	11q23.3	1	_	_	_	1	_	2	1	_	5
17	SEPT5	_	1	_	1	2	_	_	1	_	5
18	ABI1	—	2		—	2		—	_		4
19	KNL1	_	—	—		2	_	_	—	2	4
20	MAML2	—			2		—	1	—	1	4
21	MYO1F	_	3			1	_	_	_	—	4
22	PICALM	1	1	—	1	—	_	_	1	_	4
23	INRC18	1			2			1	_		4
24	FLINA	1	2	_		1	_	_	_	_	3
25	NEDL	I	I			1		1			2
20	$\Delta FEA/\Delta F5$	2	_	_	_	· _	_	_	_	_	2
28	RTRD18	2	_	_	_	_	_	_	_		2
29	CBI	<u> </u>	_	_	1	_	_	_	1	_	2
30	CEP170B	_	_	_	_	_	_	_	2		2
31	CREBBP	_	_	_			2	_	_	_	2
32	DCP1A	1	_	_	_	1	_	_	_		2
33	FOXO3A	—			2			—	_		2
34	KIAS1524	_	2	_	—	_	_	_	_	_	2
35	SEPT11		_	_	1	1	_	—	—		2
36	ABI2	—	2	_		_	_	_	_	—	2
37	ACACA		—	—	_	1	_	—	—	_	1
38	ACERI	I	_	_	_	_	_	_	1	_	1
39	ARAP 13		_	_	_	_	_	_	1		1
40	ARHGAP26	_	_	_	_	1	_	_	_	_	1
42	ARHGEF12	_	_	_			_	_	1	_	1
43	ARHGEF17	_	_	_	_	1	_	_	_		1
44	BCL9L	_	_	_	1	_	_	_	_	_	1
45	BUD13	_	_	_	_	1	_	—	_		1
46	C2CD3	_	_	—	_	—	_	_	1	_	1
47	CASP8AP2	—		—		—	—		1		1
48	CEP164	—	_	_		_	_	1	_	—	1
49	CLIA	_						1	_		1
50	CLIC	_	_	_	_	I	1	_	_	_	1
51	C14552 DCPS						I		1		1
53	FFFSFC	1	_	_	_	_	_	_			1
54	ENRP1		1	_	_	_	_	_	_	_	1
55	GAS7	_	1	_	_	_	_	_	_	_	1
56	GIGYF2	_					_		_		1
57	GMPS	_	_	_	_	_	_	_	1	_	1
58	KIF2A	_	_	_	_	_	_	1	_		1
59	LAMC3	_	—	—	—	1		—	_	—	1
60	LOC100131626	—	_		_	_	_	_	_	1	1
61	ME2		—	—	—			—	1		1
62	MKL1	—	—		1	—	—	—	_		1
63	MYH11	—	—	_	—			—	1	—	1
64	NUX4	—	1	_	_	_	_	_	1	—	1
66	NKIP3 NI ID153		I		1						1
67	PDS54	_	_	_		_	_	_	1	_	1
68	PFDN4				1		_		· 		1
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Table 1.	(Continued)										
#	Direct TPG	Infant			Pediatric			Adult			
		ALL	AML	Other	ALL	AML	Other	ALL	AML	Other	
69	PRPF19	1		_							1
70	PRRC1	_	_	_	_	_		_	1	_	1
71	RABGAP1	_	_	_	_	_	1	_	_	_	1
72	RUNDC3B	_	_	_	1	_		_	_	_	1
73	SEPT2	_	_	_	_	1		_	_	_	1
74	SMAP1	_	_	_	_	_	_	_	1	_	1
75	TCF12	_	_	_	_	_	_	_	1	_	1
76	ТОРЗА	_	_	_	_	_		_	1	_	1
77	VAV1	_	—	—	_	1		_	—	_	1
78	1p13.1	_	_	_	_	_	_	1	_	_	1
79	6q27	_	_	_	_	1	_	_	_	_	1
80	9p13.3	1	_	_	_	_	_	_	_	_	1
81	11q24.3	_	1	_	_	_	_	_	_	_	1
82	21q22	_	_	_	1	_	_	_	_	_	1
83	MLL internal inv	_	_	_	_	1	_	_	_	_	1
	SUM	692	160	24	313	339	19	415	373	10	2345

Abbreviations: AML, acute lymphoblastic leukemia; AML, acute myeloid leukemia; DCAL, Diagnostic Center of Acute Leukemia; TPG, translocation partner gene. All fusion genes that have been analyzed at the DCAL and their distribution between infant, pediatric and adult leukemia patients are shown. Total numbers are given for each patient group separated in ALL, AML and other diseases. The most frequent fusion partner genes were separated from the other genes that have been isolated less frequently by a line. Genes marked in bold represent out-of-frame *MLL–X* fusions.



**Figure 1.** Age distribution of investigated patients. The age distribution of all analyzed patients (n = 2345) is summarized. Upper part: diagram displaying ALL and AML patients. Age at diagnosis was divided into infants (0-1 year), pediatric (1-18 years) and adult patients (> 18 years). The number of ALL, AML and other patients is listed below. We also added the information about therapy-induced leukemia (TIL) patients, the number of complex *MLL* rearrangements (CL) and specified the 'non-ALL' and 'non-AML' patients (MLL, MDS, lymphoma and other) in more detail for each age group. The precise number of all patient cases is summarized on the right.

Pediatric ALL (*n* = 313) displayed 139 *AF4*, 56 *AF9*, 56 *ENL*, 12 *AF10*, 16 *AF6* and 6 *EPS15* gene fusions. Another 31 *MLL* rearrangements were characterized (*21q22*, *AF17*, *LAF4*, *BCL9L*, *CBL*, *FOXO3A*, *MAML2*, *MKL1*, 5 cases (NA) with no der(11) and only a reciprocal fusion allele, *NUP153*, *PFDN4*, *PICALM*, *RUNDC3B*, *SEPT5*, *SEPT11*, *TET1* and *TNRC18*).

Pediatric AML (n = 339) displayed 3 AF4, 132 AF9, 21 ENL, 66 AF10, 24 ELL, 6 MLL PTDs, 28 AF6 and 5 EPS15 gene fusions. Another 54 MLL rearrangements have been diagnosed (6q27, 11q23.3, ABI1, ACACA, ACTN4, AF1Q, AF17, ARHGAP26, ARHGEF17, *BUD13*, *CLTC*, *DCP1A*, *FLNA*, *KLN1*, *LAMC3*, an *MLL* gene-internal deletion, *MYO1F*, 5 cases (NA) with no der(11) and only a reciprocal fusion allele, *NEBL*, *SEPT2*, *SEPT5*, *SEPT6*, *SEPT9*, *SEPT11*, *TET1* and *VAV1*).

Adult ALL (*n* = 415) displayed 332 *AF4*, 9 *AF9*, 50 *ENL*, 1 *AF10*, 1 *ELL*, 1 *MLL* PTD, 9 *AF6* and 4 *EPS15* gene fusions. Additional 8 *MLL* rearrangements were identified (*11q23*, *ACTN4*, *CEP164*, *KIF2A*, *MAML2*, *PRRC1*, *PTD* and *TET1*).

Adult AML (*n* = 373) displayed 3 *AF4*, 90 *AF9*, 14 *ENL*, 33 *AF10*, 45 *ELL*, 98 *MLL* PTDs, 38 *AF6* and 5 *EPS15* gene fusions. Another 47



**Figure 2.** Classification of patients according to age classes and disease type. Top: frequency of most frequent TPGs in the investigated patient cohort of MLL-r acute leukemia patients (n = 2345). This patient cohort was divided into ALL (left) and AML patients (right). Gene names are written in black, percentages are indicated as white numbers. Fifty-three patients could not be classified into the ALL or the AML disease types, respectively. Middle: TPG frequencies for the infant, pediatric and adult patient group. Bottom: subdivision of all three age groups into ALL and AML patients. Negative numbers refer again to the number of patients that were neither classified to the 'ALL' nor to the 'AML' subgroup.

*MLL* rearrangements were detected (*1p13.1*, *11q23*, *AF1Q*, *AKAP13*, *AP2S2*, *ARHGEF12*, *C2CD3*, *CASP8AP2*, *CBL*, *CEP170B*, *DCPS*, *GMPS*, *ME2*, *AF17*, *MYH11*, 2 cases (NA) with no der(11) and only a reciprocal fusion allele, *NOX4*, *PDS5A*, *PICALM*, *SEPT5*, *SEPT6*, *SEPT9*, *SMAP1*, *TCF12*, *TET1* and *TOP3A*).

On the basis of the above distribution, about 96% of all ALL patients (n = 1420) were characterized by the fusion genes MLL-AF4 (~57%), MLL-ENL (~18%), MLL-AF9 (~13%), MLL-AF10 (~4%), MLL-EPS15 (~2%) and MLL-AF6 (~2%). About 83% of all AML patients (n = 872) were characterized by the fusion genes MLL-AF9 (~30%), MLL-AF10 (~16%), MLL-ELL (~11%), MLL PTDs (~12%), MLL-AF6 (~8%), MLL-ENL (~4%) and MLL-SEPT6 (~2%). These results are in line with recently published data about the frequency and distribution of different MLL fusion partner genes.<sup>16–18</sup> This updated information is quite important for diagnosis and has already been used to establish a fast reversetranscription PCR-based multiplex screening method.<sup>19</sup> Additional information about the distribution of translocation partner genes (TPGs) in major disease subgroups (different B-cell developmental stages, T-ALL or French-American-British (FAB) M0-M7 for AML patients) have been summarized in Supplementary Figures S1 and S2. Here the different FAB classes in AML show a guite distinct pattern of fusion genes with some of the major fusion partners missing in distinct FAB groups M0-M7. For example, in FAB M0 AF4 and ELL are missing, in FAB M1 ENL is missing, in FAB M2 AF4, AF10 and AF6 are missing and so on. In FAB class M6 and M7, only certain fusion genes could be identified. In B-ALL stages, no such exclusion patterns were observed, rather a shift for specific fusion genes, while T-ALL is mainly composed of ENL and AF6 gene fusions (see also below).

Breakpoint distribution according to clinical subtypes

We also investigated the distribution of chromosomal breakpoints within the *MLL* breakpoint cluster region in all investigated clinical subgroups. Briefly, the major breakpoint cluster region is localizing between *MLL* exon 9 and *MLL* intron 11, where the majority of patients (93.5%) had their individual breakpoints (n = 2192). Only 153 patients (6.5%) had their breakpoint outside of the major breakpoint cluster region (Supplementary Figures S3–S5 and Supplementary Table S6).

As the localization of breakpoints may have an impact on cancer biology and clinical behavior, we started to analyze the breakpoint distribution for all clinical subgroups and compared them with the 'mean distribution' (MD) observed for all 2345 patients. We decided not to use a 'random distribution model' of chromosomal breakpoints, because this is only based on the length of each DNA region. However specific features in MLL intron 9 (four Alu repetitive elements of which three are transcriptionally active) and MLL intron 11 (sensitivity against cytotoxic drugs, a DNase1 hypersensitive site,<sup>20</sup> an apoptotic cleavage site,<sup>21</sup> an RNA polymerase II-binding site<sup>22</sup> and topoisomerase II-binding sites<sup>23</sup>) may account for a specific increase of DNA double-stranded breaks due to specific molecular features of the chromatin, or, breakpoints differ because of a selection process for resulting MLL fusion proteins.

For our analyses, we subdivided the *MLL* breakpoint cluster region into three subregions: (A) exon 9–intron 9=1761 bp; (B) exon 10–intron 10=679 bp; (C) exon 11–intron 11–exon 12–intron 12 and exon 13=5026 bp. The observed 'MD' for these three *MLL* breakpoint regions was A=37.0%, B=19.8% and C=40.1% for all 2345 patients listed in Supplementary Table S7. In these analyses,

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#	Cytogenetic abnormality	Breakpoint	TPG	References	Leukemia type
A. ML	L fusion in-frame				
	t(1;11)(p32;q23)	1p32	EPS15/AF1P	Bernard et al. (1994)	ALL, BAL, AML, CML
	t(1;11)(q21;q23)	1q21	MLLT11/AF1Q	Tse <i>et al.</i> (1995)	AML, t-AML, ALL, t-ALL, BAL
	t(2;11)(p23.3;q23)	2q23.3	ASXL2	Haferlach <i>et al.</i> (2016)	t-AML
	ins(2;11)(q11.2;q23)	2q11.2	AFF3/LAF4	von Bergh <i>et al</i> . (2003)	ALL
	t(2;11)(q33;q23)	2q33	ABI2	Coenen et al. (2012)	AML
	t(2;11)(q37;q23)	2q37	SEPT2	Cerveira et al. (2006)	t-AML, AML, t-MDS
	t(2;11)(q37.1;q23)	2q37.1	GIGYF2	Not published yet	ALL
	t(3;11)(p21;q23)	3p21	NCKIPSD/AF3P21	Sano <i>et al</i> . (2000)	t-AML
	t(3;11)(p21.1;q23)	3p21.1	DCP1A	Meyer et al. (2008)	ALL, AML
)	t(3;11)(q13.13;q23)	3q13.13	KIAS1524	Coenen et al. (2011)	AML
	t(3;11)(q24;q23)	3q24	GMPS	Pegram <i>et al</i> . (2000)	t-AML, t-MDS
	t(3;11)(q28;q23)	3q28	LPP	Daheron <i>et al.</i> (2001)	t-AML
	t(4;11)(p14;q23)	4p14	PDS5A	Meyer et al. (2011)	t-AML, AML
	t(4;11)(p11;q23)	4p11	FRYL	Hayette <i>et al</i> . (2006)	t-ALL, t-AML, t-MDS
	t(4;11)(q21.1;q23)	4q21.1	SEPT11/FLJ10849	Kojima <i>et al</i> . (2004)	T-ALL, CML, t-ALL, t-AML
	t(4;11)(q21;q23)	4q21	AF4/AFF1	Gu et al. (1992)	ALL, t-ALL, BAL, AML
'	t(4;11)(q35.1;q23)	4q35.1	SORBS2/ARGBP2	Pession <i>et al.</i> (2006)	AML
	t(5;11)(q12.1;q23)	5q12.1	KIF2A	Not published yet	ALL
	complex abnormalities	5q12.3	CENPK/FKSG14	Taki et al. (1996)	AML
	t(5;11)(q23.2;q23)	5q23.2	PRRC1	Douet-Guilbert et al. (2014)	t-ALL
	ins(5;11)(q31;q13q23)	5q31	AFF4/AF5Q31	Taki et al. (1999)	ALL
	t(5;11)(q31;q23)	5q31	ARHGAP26/GRAF	Borkhardt et al. (2000)	JMML
	t(6;11)(q13;q23)	6q13	SMAP1	13	AML
	t(6:11)(g15:g23)	6q15	CASP8AP2	Park et al. (2009)	AML
	$t(6.11)(q_{2}1.q_{2}3)$	6q21	EOXO3/AE6O21	Hillion et al. (1997)	t-AMI t-ALI
	$t(6,11)(q_27,q_23)$	6q27	MITTA/AF6	Prasad et al. (1993)	T-ALL AML T-AML ALL
	t(7,11)(n22,1,n23)	7n221	TNRC18/KIAS1856	Mover et al. $(7008)$	
	t(7,11)(p22,1,q23) t(7,11)(a11,23,a23)	7 p22.1 7 a 11 23	CLIDS	Not published yet	
	(7,11)(q11.23,q23) +(7,11)(q11.23,q23)	7011.25		16	
	(7,11)(q21.12,q23) +(7,11)(q221.q23)	7q21.12 7a22.1	FUNC	Hafarlach at al (2016)	1-ALL + AMI
	$t(7,11)(q_{3}2,1,q_{2}3)$	7432.1 0p12	FLINC CLTA	Not published yet	
	t(9;11)(p13;q23)	9p13		Not published yet	
	l(9;11)(p22;q25)	9pzz	IVILLIS/AFY	Nakamura et al. (1995)	AIVIL, L-AIVIL, ALL, I-ALL, DAL
	t(9;11)(q33.2;q23)	9q33.2	DABZIP/AF9Q34	Von Bergn <i>et al.</i> (2004)	AML
	$t(9;11)(q_{34},11;q_{23})$	9q34.11	KABGAPT	Not published yet	ALL
	Ins(11;9)(q23;q34)Inv(11)(q13)(q23)	9q34	FINBP I/FBP I /	Fuchs et al. (2001)	AML
	t(9;11)(q34.12;q23)	9q34.12	LAMC3		t-AML
	ins(10;11)(p12;q23)	10p12	NEBL	Coser et al. (2010)	AML
	ins(10;11)(p12;q23q13)	10p12	MLLT10/AF10	Chaplin <i>et al.</i> (1995)	AML, t-AML, ALL, I-ALL, BAL
	t(10;11)(p11.2;q23)	10p11.2	ABI1	Taki <i>et al.</i> (1998)	AML
	t(10;11)(q21;q23)	10q21	TET1/LCX	Ono <i>et al</i> . (2002)	AML, ALL
	inv(11)(p15.3q23)	11p15.3	NRIP3	5	AML
	inv(11)(q12.1q23)	11q12.1	BTBD18	Alonso et al. (2012)	ALL
	inv(11)(q12.2q23)	11q12.2	PRPF19	16	ALL
	t(11;11)(q13.4;q23)	11q13.4	ARHGEF17	Teuffel et al. (2005)	AML
	inv(11)(q13.4q23)	11q13.4	C2CD3	15	AML
	inv(11)(q14q23)	11q14	PICALM/CALM	Wechsler et al. (2003)	AML, ALL
	inv(11)(q21q23)	11q21	MAML2	14	T-ALL,t-T-ALL, t-AML, t-MDS
	del(11)(q23q23.3)	11q23.3	CBL	Fu <i>et al.</i> (2003)	AML, t-AML, ALL
	del(11)(q23q23.3)	11q23.3	ARHGEF12/LARG	Kourlas <i>et al.</i> (2000)	AML, t-AML
	del(11) )(q23q24.2)	11q24.2	DCPS	13	AML
	t(11;12)(q23;p11.23)	12p11.23	ITPR2	Haferlach <i>et al</i> . (2016)	t-MDS
	t(11;12)(q23;q13.2)	12q13.2	SARNP/CIP29	Hashii <i>et al.</i> (2004)	AML
	t(11;14)(q23;q23.3)	14a23.3	GPHN	Kuwada <i>et al.</i> (2001)	AML, t-AML
	t(11;14)(q23;q32.33)	14q32.33	CEP170B/KIAA0284	Burmeister et al. (2008)	t-AML
	t(11:15)(q23:q14)	15a14	KNL1/CASC5	Havette et al. (2000)	AML, t-MDS, ALL,
	t(11:15)(a23:a15.1)	15q15.1	ZFYVE19/MPFYVE	Chinwalla <i>et al.</i> (2003)	AML
	t(11:15)(a23:a21)	15g21	TCF12	Not published vet	t-AMI
	t(11:15)(a23:a25.3)	15a253	AKAP13	16	t-AMI
	t(11.16)(a23.n13.3)	16n133	CRERRP/CRP	Taki et al (1997)	t-MDS t-AMI AMI t-AII t-
	t(11.16)(q23.p13.3)	16p13.5	MYH11	16	
	(11,10)((42),(13,11)) t(11,17)(a)(a)(a)(13,11)	10p13.11 17p12 1	GAS7	Maganias at al (2000)	
	(11,17)(423,173,17) +(11,17)(a22,171,2)	17p13.1		Horbaux at $a^{1}$ (2012)	
	t(11,17)(q23;p11.2)	17p11.2	I UPSA	$\begin{array}{c} \text{Herbulk et al. (2012)} \\ \text{Strobl at al. (2002)} \end{array}$	
	l(11;17)(q23;q12)	17012	LASPI	13	
	lns(11;1/)(q23;q21)	1/q21	ACACA		AML
	t(11;17)(q23;q21)	1/q21	MLL16/AF17	Prasad et al. (1994)	AML, ALL
	t(11;17)(q23;q23.1)	17q23.1	CLTC	Not published yet	AML
	t(11;17)(q23;q25)	17q25	SEPT9/AF17Q25	Usaka et al. (1999)	t-AML, AML, MDS, ALL
2	t(11·18)(a23·a21)	18a21	ME2	Szotkowski et al. (2015)	t-AML

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able 2	. (Continued)				
#	Cytogenetic abnormality	Breakpoint	TPG	References	Leukemia type
59	t(11;19)(q23;p13.3)	19p13.3	MLLT1/ENL	Tkachuk et al. (1992)	ALL, T-ALL, AML, BAL, t-AL
70	t(11;19)(q23;p13.3)	19p13.3	ACER1/ASAH3	Lo Nigro <i>et al.</i> (2002)	ALL
71	t(11;19)(q23;p13.3)	19p13.3	SH3GL1/EEN	So et al. (1997)	AML
2	ins(11;19)(q23;p13.3)	19p13.3	VAV1	15	AML
'3	t(11;19)(q23;p13.2)	19p13.2	MYO1F	Lo Nigro <i>et al.</i> (2002)	AML
'4	t(11;19)(q23;p13.1)	19p13.1	ELL	Thirman <i>et al.</i> (1994)	ALL, BAL, AML, t-AML
5	t(11;19)(q23;q13)	19q13	ACTN4	Burmeister et al. (2009)	t-ALL, t-AML
6	t(11;20)(q23;q11.21)	20q11.21	MAPRE1	Fu et al. (2005)	ALL
7	t(11;20)(q23;q13.2)	20q13.2	PFDN4	Not published yet	T-ALL
8	t(11;22)(q23;q11.21)	22q11.21	SEPT5/CDCREL	Megonigal <i>et al.</i> (1998)	AML, T-ALL
9	t(11;22)(q23;q13)	22q13	MKL1	Not published yet	ALL
0	t(11;22)(q23;q13.2)	22q13.2	EP300/P300	Ida et al. (1997)	t-AML
	t(X;11)(q13.1;q23)	Xq13.1	FOXO4/AFX	Parry et al. (1994)	I-ALL, ALL, t-ALL,CLL, AML
32	ins(X;11)(q24;q23)	Xq24	SEP16	Borkhardt et al. (2001)	AML
33	lns(X; II)(q26.3; q23)	Xq26.3	C14552	Cerveira et al. (2010)	BAL
4	Ins(11;X)(q23q28q13.1)	Xq28	FLNA	De Braekeleer et al. (2009)	AML
. ML	t(3;11)(p21.3;q23)	3p21.3	SACM1L	34	N/A
	t(3:11)(g21.3:g23)	3a21.3	EEFSEC/SELB	13	ALL
	t(6:11)(p22.3:q23)	6p22.3	NUP153	Not published vet	ALL
	inv(11)(p15.5q23)	11p15.5	AP2S2	16	AML
	complex	11a14	NOX4	Not published vet	ALL
,	t(11:15)(q23.3:q21)	11a23.3	LOC100131626	33	MDS
	inv(11)(g23.3g23)	11g23.3	BUD13	16	AML
3	del(11)(g23g23.3)	11q23.3	CEP164	16	t-ALL
)	del(11)(g23g23.3)	11q23.3	BCL9L	14	ALL
0	t(2;11;19)(p23.3;q23;p13.3)	19p13.3	LOC100128568	15	AML
No	partner gene fused to 5'-MLL gene				
	t(1:11)(p13.1:q23)	1,51a1		16	PMF
	t(6:11)(g27:g23)	6q27		Not published vet	AML
	t(9:11)(p13.3:q23)	9p13.3		16	t-ALL
	t(11:11)(q23:q23.3)	11a23.3		16	ALL, AML
,	t(11;11)(q23;q24.3)	11q24.3		16	AML
6	t(11;21)(q23;q22)	21q22		16	t-ALL
D. No 1 2 3 4 5	t characterized at the molecular level ( t(1;11)(p36;q23) t(1;11)(q31;q23) t(1;11)(q32;q23) t(2;11)(p21;q23) t(2;11)(q37;q23)	published by oth	ers)		
	t(3;11)(p13;q23)				
,	t(4;11)(p11;q23)				
	t(6;11)(q13;q23)				
)	t(7;11)(p15;q23)				
0	t(7;11)(q22;q23)				
1	t(7;11)(q32;q23)				
2	t(8;11)(q11;q23)				
3	t(8;11)(q21;q23)				
4	t(8;11)(q24;q23)				
5	t(9;11)(p11;q23)				
6	t(9;11)(q33;q23)				
7	t(10;11)(q25;q23)				
8	t(11;11)(q11;q23)				
9	t(11;11)(q13;q23)				
C	t(11;11)(q21;q23)				
1	t(11;12)(q23;p13)				
2	t(11;12)(q23;q13)				
3	t(11;12)(q23;q24)				
4	t(4;13;11)(q21;q34;q23)				
5	t(11;14)(q23;q11)				
6	t(11:14)(g23;g32)				
7	t(11:15)q23:q15)				
8	t(11:17)(a23:a11)				
9	t(11:17)(q23;q23)				
0	t(11:18)(a23:a12)				
1	t(11:18)(q23;q23)				
	t(11·20)(a23·a13)				
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Table	2. (Continued)				
#	Cytogenetic abnormality	Breakpoint	TPG	References	Leukemia type
33 34 35	t(11;21)(q23;q11) t(Y;11)(p11;q23) t(X;11)(q22;q23)				
Abbre Diagr relate gene ident	eviations: AML, acute lymphoblastic nostic Center of Acute Leukemia; JM d AML; t-MDS, therapy-related MDS; name, the appropriate reference anc ified at the DCAL during the last de	leukemia; AML, acute IML, juvenile myelom TPG, translocation pa d observed disease ty cade. In addition, 6 c	e myeloid leuke onocytic leuke artner gene. Lis pe. Genes mark loned gene loc	emia; BAL, bilineal acute leukem mia; MLL, mixed lineage leuken t of the cytogenetic localization ted as 'not published yet' are cor i and 35 cytogenetic chromoso	ia; CML, chronic myelogenous leukemia; DCAL, nia; t-ALL, therapy-related ALL; t-AML, therapy- of all yet-characterized direct TPGs ( $n = 94$ ), the mpletely new. All references in italics have been me loci have been identified.

all patients were investigated for their fusion partner gene in correlation with age at diagnosis, gender, patient group, therapyinduced leukemia, complex genetic rearrangements, origin of patient and breakpoint distribution. Here a significant deviation from the 'MD' was observed for AF1Q, AF6, AF10, ENL, EPS15, SEPT6, SEPT9, AF17 and MLL PTDs. The fusion partner genes ENL and SEPT6 had preferentially MLL intron 11 breaks, while all others tend to bear MLL intron 9 recombination events. Of interest, also therapy-induced acute leukemias differ significantly in their 'MD', with a tendency for MLL intron 11 breaks.

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A detailed analysis for the most frequent MLL fusion partner genes is depicted in Supplementary Table S8. Here we separated according to fusion partners and patient subgroup (infant I, pediatric P and adult A) with regard to several other parameters such as age, gender, therapy-induced, complex translocation, origin and disease type. The overall breakpoint distribution of all seven most frequent genetic aberrations with more than 2000 patients was not significantly deviating from the MD of all patients. However, significant changes were observed for patient subgroups bearing chromosomal translocations to AF4 (I and A), AF9 (A), ENL (I and P), AF10 (P), ELL (I, P and A), AF6 (I, P and A) and MLL PTD patients (P and A). This clearly demonstrates that certain fusion genes have a selective preference for distinct breakpoints, most likely because of the resulting function of a given fusion protein. As an example, AF6 fusions in ALL and AML patients are mostly occurring in MLL intron 9 (or even upstream), while infant AF4 and infant/pediatric ENL fusions tend to have breakpoints within MLL intron 11. Similar observations were made for the more rare fusion partner AF1Q (significantly toward MLL intron 9), AF17 (significantly toward MLL introns 8 and 9), EPS15 (significantly toward MLL intron 11 in adult patients), SEPT6 (significantly toward MLL intron 11 in pediatric and adult patients) and SEPT9 (significantly toward MLL introns 7-9).

To evaluate these data further, we correlated the breakpoint distribution with the age of patients. We have done so for ALL and AML patients (Supplementary Figures S9 and S10). These analyses revealed that the disease subtypes (ALL and AML) divide patients into two groups (ALL more in MLL intron 11 breakpoints; AML more in MLL intron 9 breakpoints). However, these breakpoint tendencies seem to change with age. Thus, young patients usually display MLL intron 11 breakage, while older patients have more breaks in MLL intron 9. This is true for all investigated subgroups (AF4, ENL and AF9) where we had enough patients to perform this type of analysis and to obtain a significant result. Vice versa, young AML patients usually prefer MLL intron 9 breakage, while older patients have more breaks in MLL intron 11. This has been done also for the AF9, AF10 and ELL subgroups. The cross-over points were 10-14 years in ALL patients and 75 years in AML patients. MLL-ELL patients within the AML group are somehow different from all other subgroups because they start very early on with a preference for MLL intron 11 (all patients above 1 year of age) and display no cross-over point. These breakpoint preferences and their change with age are indicating that two different molecular mechanisms are driving MLL rearrangements: one is a transcription-coupled hot spot that is quite sensible for external cytotoxic triggers (*MLL* intron 11), while the other is presumably based on transcriptionally active ALU repeats where POL III and POL II transcription is causing torsional DNA stress.

Another important point is the outcome of patients that is linked to the distribution of chromosomal breakpoints within the MLL breakpoint cluster region.<sup>24</sup> Basically, the outcome of leukemia patients with breakpoints in MLL intron 11 was worse compared to those patients with upstream breakpoints. A rational explanation for this observation came from functional studies of the plant homeodomain (PHD) domain of the MLL protein, encoded by MLL exons 11–16 (Supplementary Figure S11). This domain is built up by PHD1, PHD2 and an enhanced PHD3. The first three PHD domains are separated by the adjacent bromodomain and another enhanced PHD4. PHD3 has an important dual function, because it either binds to the CYP33/PPIE protein<sup>25,26</sup> or to methylated lysine-4 residues of histone H3.<sup>27</sup> Binding of PHD3 to H3K4<sub>me2/3</sub> peptides is greatly enhanced by the adjacent bromodomain,<sup>28</sup> but binding of the prolyl-peptidyl isomerase CYP33/PPIE confers a cis-trans isomerization of proline-1665. This enables binding to BMI1 and associated repressor proteins (HDAC/CBX4/KDM5B). This molecular switch determines the human MLL protein of being a transcriptional activator/maintenance factor or a transcriptional repressor. Noteworthy, PHD2 and PHD3 also bind to E3-ligases (CDC34 and ASB2, respectively), which control the steady-state stability of the MLL protein.<sup>29,30</sup> As shown recently by our group, breakpoints within MLL intron 11 destroy the dimerization capacity of the PHD1-3 domain.<sup>31</sup> This also disables binding to the BMI1 repressor complex.<sup>32</sup> Thus, a breakpoint within MLL intron 11 has functional consequences for the resulting fusion proteins, which may explain the clinical finding of the altered outcome.<sup>24</sup>

## The MLL recombinome

On the basis of the results obtained in the present and previous studies, 13-16 a total of 84 direct TPGs and their specific breakpoint regions have now been identified, all of which generate an inframe MLL fusion protein (Table 2A). Additional 10 MLL fusions were characterized that were fused out of frame to translocation partner genes (Table 2B). In the latter cases, alternative splicing may still allow to generate an MLL-fusion protein, however, this was not investigated here. Another 6 loci have been cloned where the 5'-portion of MLL was not fused to another gene (Table 2C). 3'-Race and reverse-transcription PCR experiments with several exon combinations were performed to identify potential fusion transcripts. But no in-frame fusion RNAs could be identified. Therefore, these 16 unusual MLL rearrangements—where neither any dimerization nor a transcriptional activation domain is present probably represent a subclass of MLL abnormalities, which have per se no or only a weak ability to transform hematopoietic cells and are only identified in the context of other genetic abnormalities in hematopoietic stem cells.33,34

In 31 additional cases we were not able to identify a der(11) fusion gene at all. This could be either attributed to a technical

problem (for example, a too long genomic fragment) or to the fact that no der(11) exists in these few patients. However, in 22/31 cases we successfully identified a reciprocal MLL fusion allele, while 9 cases carried no detectable direct or reciprocal fusion gene. This subgroup (n=31) was included in the group of 'complex *MLL* rearrangements' (n = 247) because of the extending class of 'reciprocal MLL fusion genes' (Supplementary Table S12). Within this group of patients with complex *MLL* rearrangements, a total of 32 reciprocal MLL fusions represent in-frame fusions (marked in red in Supplementary Table S12), while 215 fusions were out-of-frame fusions at the genomic DNA level (88 gene loci/ 127 partner genes; shown in black in Supplementary Table S12). Finally, there were still 35 chromosomal translocations of the human MLL gene that were characterized in the past by cytogenetic methods, but that were never analyzed at the molecular level (Table 2D). Thus, the MLL recombinome to date comprises 94 different 'direct TPGs' (decoding the MLL N terminus) and 6 different 5'-MLL genes fused only to genomic DNA. On the other hand we have now 247 'reciprocal TPGs' (bearing the MLL C terminus) that are deriving from complex rearrangements with already-known 'direct TPGs'. It is noteworthy that nearly all of these reciprocal MLL fusion (243 out of 247) are per se able to express only the 3'-MLL portion, named MLL\*, due to a gene internal promoter located upstream of MLL exon 12.22

# Novel translocation partner genes

Apart from the many new *MLL* fusion genes that have already been discovered at the DCAL and published in the last decade (Tables 2a–c; *n* = 40), we present additional 11 novel TPGs (marked as 'not published yet'): *GIGYF2* (GRB10-interacting GYF; 2q37.1; 1299 aa); *KIF2A* (kinesin heavy chain member 2A; 5q12.1; 706 aa); *CLIP2* (CAP-GLY domain-containing linker protein 2; 7q11.23; 1046 amino acids (aa)); *CLTA* (clathrin, light chain A; 9p13.3; 248 aa); *RABGAP1* (RAB GTPase-activating protein 1; 9q33.2; 1069 aa); *TCF12* (transcription factor 12; 15q21.3; 682 aa); *CLTC* (clathrin, heavy chain; 17q23.1; 1,675 aa); *PFDN4* (prefoldin subunit 4; 20q13.2; 134 aa); *MKL1* (megakaryoblastic leukemia (translocation) 1; 22q13.1; 931 aa); *NUP153* (nucleoporin 153 kDa; 6p22.3; 1475 aa); and *NOX4* (NADPH oxidase 4; 11q14.3; 578 aa).

The *Drosophila* GIGYF2 protein ortholog was shown to be a modulator of autophagy that controls neuron and muscle homeostasis.<sup>35</sup> GIGYF2 binds directly to AGO2 and is involved in siRNA-mediated post-transcriptional silencing.<sup>36</sup> A quite specific feature of GIGYF2 is to build a complex together with eIF4E and ZNF598 to selectively block the process of translation of distinct capped mRNAs.<sup>37</sup> Several papers have linked *GIGYF2* also to Parkinson's disease, however, these data are so far not sufficiently significant.

KIF2A is a member of the kinesin-13 family and involved in spindle assembly at the metaphase I-anaphase I transition of oocytes.<sup>38,39</sup> Moreover, genetic mutations in the motor domain of this protein is associated with cortical malformation syndromes such as microcephaly.<sup>40</sup> Vice versa, overexpression of KIF2A has been diagnosed in different cancers, because KIF2A expression and phosphorylation influences microtubuli dynamics, which is important for tumor cell migration and metastasis.<sup>41</sup>

*CLIP2* has been discovered as overexpressed biomarker after radiation in papillary thyroid carcinomas, usually accompanied by a gain of chromosome band 7q11.<sup>42</sup> This disease has been frequently diagnosed as the main consequence of the Chernobyl accident.

CLTA, also named clathrin light chain A, is involved in vesicle trafficking and endocytosis. However, a recent paper demonstrated that CLTA has a role on the migration of tumor cells.<sup>43</sup> This is in part due to the fact that CLTA interacts with Huntingtininteracting protein, involved in the regulation of the actin cytoskeleton. Upon depletion of clathrin light chains a steadystate downregulation of  $\beta$ 1-integrins was observed because of defects in vesicle recycling.

RABGAP1 is highly specific for RAB6A and has a role in microtubule nucleation at the centrosome. It also participates in a RAB6A-mediated pathway involved in the metaphase—anaphase transition (Mad2-spindle checkpoint).<sup>44</sup>

TCF12, also known as HeLa E-box binding protein, is quite interesting as it controls the osteogenic differentiation of mesenchymal stem cells in the bone marrow.<sup>45</sup> This basic helix–loop–helix transcription factor was also found to be mutated in anaplastic oligodendroglioma.<sup>46</sup> TCF12 is able to bind to TWIST1 and involved in the early differentiation pathways of thymic T cells (DN3-> DN4 and ISP-> DP).<sup>47</sup>

*CLTC* has been identified in complex chromosomal rearrangements causing the expression of the *CLTC–ALK* fusion in blastic plasmacytoid dendritic cell neoplasm.<sup>48</sup> A more recent work has found *CLTC* mutations are associated with neuronal malformations and intellectual developmental delays.<sup>49</sup> This is due to the fact that clathrin heavy and light chains (CLTA and CLTC) are involved in vesicle trafficking, vesicle recyling and neurotransmitter release.

PFDN4 has been linked to colorectal cancer, however, inversely correlated with outcome (low expression has poorer outcome). A knockdown of this gene was correlated with increased cell growth and invasiveness.<sup>50</sup>

MKL1 interacts with the transcription factor myocardin, a key regulator of smooth muscle cell differentiation. The encoded protein is predominantly nuclear and may help transduce signals from the cytoskeleton to the nucleus. This gene is involved in a specific translocation event that creates a fusion of this gene and the RNA-binding motif protein-15 gene. This specific t(1;22)(p13; q13) translocation has been associated with the development of acute megakaryocytic leukemia.<sup>51</sup>

NUP153 s a highly versatile protein, involved in nuclear pore functions, pore architecture, nuclear import and export, *de novo* pore formation after mitosis and destruction of NUP153 during apoptosis.<sup>52</sup>

NOX4 is NADPH oxidase 4 that is important in the regulation of glycolysis and glutamate metabolism. Disruption of *NOX4* by CRISPR/Cas9 is inhibiting cell growth of HeLa cells, indicating that *NOX4* is quite important as metabolic regulator in tumor cells.<sup>53</sup> *NOX4* has been identified in many tumors as a relevant gene.

## T-ALL cases

A tiny fraction of investigated patients were diagnosed with a T-ALL (n = 59) (Supplementary Figure S13). This group of patients is mainly characterized by *MLL* fusion with *AF6* (n = 23) and *ENL* (n = 22). Other fusions were AF4, AF9, AF10, MAML2, PFDN4, RUNDC3B, SEPT5, SEPT11, TNRC18 and 1 reciprocal USP20-MLL fusion. Only in the cohort of MLL-AF6 patients, quite unusual MLL breakpoints were observed (n=4), where the chromosomal breakpoint in the MLL gene was diagnosed within intron 21 and 23. This is guite important because such a far away downstream breakpoint includes the complete PHD1-3, the bromodomain as well as the complete enhanced PHD4 domain of MLL into the fusion protein with AF6 (Supplementary Figure S11). These additional 581 amino acids could be an important hint for the importance of these MLL domains in T-ALL. The PHD1-3 and bromodomain exert important regulatory functions to the MLL N terminus, such as chromatin reading, protein stability or CYP33 binding. In the latter case, binding of the BMI1 repressor complex will reverse the function of the MLL-AF6 fusion by repressing gene transcription. This is quite interesting and provides a new research aspect for MLL-AF6.

# Therapy-induced leukemia cases

We also investigated the therapy-induced patient cases (n = 110; Supplementary Table S14). The dominant partner genes are AF9 (n = 41), ELL (n = 11), AF4 (n = 11) and ENL (n = 10). All other fusions 282

(*n* = 23) have been identified one to four times. To our surprise, the AF9 cases were shifting from MLL intron 9 breaks to MLL intron 11 breaks. Some MLL fusions can only be found in therapy-induced acute leukemia and not in patients with *de novo* diseases: *ACTN4*; *AKAP13*; *ARHGEF12*; *FOXO3A*; *GMPS*; *LAMC3*; *ME2*; *PDS5A*; *PRRC1*; and *TCF12*. As expected, therapy-induced acute leukemias were only diagnosed in pediatric and adult patients, not in infants.

# Spliced fusions

Spliced fusions are rare events except for the *ENL* fusion gene (n = 302). In the latter cases, about 50% of all breakpoints localize far upstream of *ENL* exon 1 (n = 153; Supplementary Table S15). In these cases, no reciprocal fusion protein can be made, only an *MLL–ENL* fusion transcript.<sup>54</sup> For the other cases, a similar scenario was found. In all these cases, a 3'-terminal truncated *MLL* was recombined upstream of *PRPF19* (1 out of 1 case), *ELL* (8 out of 97 cases), *MYO1F* (1 out of 4 cases), *EPS15* (9 out of 38 cases), *AF4* (1 out of 839 cases), *AF6* (3 out of 95 cases) and *AF9* (2 out of 449 cases). A total of 180 cases were identified that show this unusual peculiarities.

## DISCUSSION

Here we present an update of the '*MLL* recombinome' associated with different hematologic malignancies, and in particular with acute leukemia (ALL and AML). All our analyses were performed by using small amounts of genomic DNA that were isolated from bone marrow or peripheral blood samples (n = 2345) of leukemia patients. In some cases, we analyzed cDNA from a given patient to validate the presence of *MLL* spliced fusions, or to investigate alternative splicing of RNA generated from the investigated *MLL* fusion genes. The results of this study allow to draw several conclusions.

The applied long-distance inverse PCR technique allowed to identify direct and reciprocal *MLL* fusions, *MLL* gene-internal duplications, chromosome 11 inversions, chromosomal 11 deletions and the insertion of chromosome 11 material into other chromosomes, or vice versa, the insertion of chromatin material of other chromosomes into the *MLL* gene. It is noteworthy to mention that no other technique (for example, next-generation

sequencing) displays such a high identification of chromosomal fusion sites so far. Even paired-end mRNA analysis by next-generation sequencing has a discovery rate of 60–70% only, however, RNA-based technologies do not provide the patient-specific chromosomal fusion sequences that could be used for MRD analyses. Thus, this 'old-fashioned' method is still state of the art and will be used also in the future to gain additional information of the MLL recombinome.

Moreover, we extended our knowledge on complex *MLL* rearrangements (n = 247) leading to a large collection of reciprocal *MLL* fusions (Supplementary Table S12). About 13% represent inframe fusions that can be readily expressed into reciprocal fusion proteins. All other represented out-of-frame fusions with either a chromosomal locus or a reciprocal TPG. Out-of-frame fusions such as *IKZF1–MLL*, *PBX1–MLL* or *JAK1–MLL* most likely represent a situation where such TPGs were destroyed, creating a typical loss-of-function situation. However, even those out-of-frame MLL fusions still allow to transcribe and express a 5'-truncated MLL protein, termed MLL\*.<sup>22</sup> This shorter version of MLL has no ability to bind Menin1, LEDGF or MYB, but still carries all enzymatic functions necessary to carry out H4K16 acetylations by the associated MOF protein or H3K4 methylation by the SET domain complex.

The analysis of 2345 *MLL* fusion alleles led to the discovery of 51 novel TPGs in the past 12 years, of which 40 have already been described (Tables 2a–c). Eleven TPGs are completely new and have not been published yet. Together with 49 MLL fusions that have been described by others, we can present today a total of 94 *MLL* fusions that have been characterized at the molecular level and 6 *MLL* translocations to different genetic loci (with no obvious gene fusion). All these *MLL* fusions provide a rich source for future analyses of oncogenic MLL protein variants. We have summarized all yet-known MLL fusion partner genes in Figure 3, according to their disease type/subtype in which they have been diagnosed.

According to our data, the 7 most frequent rearrangements of the *MLL* gene differ significantly in the cohorts of infant, pediatric and adult leukemia patients. We also observed significant tendencies that correlate with fusion genes, gender or age at diagnosis. As an example *MLL–AF10* (P=0.0024) occur more frequently in the male group of patients, while females were more affected by *MLL–AF4* fusions (P=0.00576). The most striking



**Figure 3.** Classification of all yet known fusion partner genes by disease. All TPGs identified were grouped by their diagnosed disease type. Genes have been diagnosed in ALL, t-ALL, t-AML, AML, T-ALL, MLL, bilineal acute leukemia (BAL), MDS, t-MDS, chronic myelogenous leukemia (CML), t-CML, juvenile myelomonocytic leukemia (JMML) and lymphoma. Genes in the intersection belong to two different groups. Bold-marked TPGsb are the most frequent ones.

finding was that breakpoint distributions differ significantly when concerning distinct TPGs and age groups. It is well-known that breakpoints in infants occur more frequently in MLL intron 11. We could validate this finding for infants with MLL-AF4, infant/ pediatric patients with MLL-ENL fusions and pediatric/adult patients with MLL-ELL fusions. However, we observed a contrary situation in adult patients with MLL-AF9 or pediatric patients with MLL-AF10 fusions. Quite surprising was the breakpoint distribution for MLL-AF6 fusions that displayed a clear preference for MLL intron 9 recombinations. Again, these deviations from the observed mean breakpoint distribution (MD) are an argument for differences in the biology of the resulting fusion proteins with respect to oligomerization or factor-binding dependency. Alternatively, it may reflect differences in the biology of transformed cell types, or, reflect different situations during the onset of these translocations (in utero exposition with poisons vs postnatal acquirement).

An important translational aspect of this study is the establishment of patient-specific DNA sequences that can be used for monitoring MRD by quantitative PCR techniques. Because of the fact that a given MLL fusion allele is genetically stable and a monoallelic marker for each tumor cell, a more reliable quantification and tracing of residual tumor cells becomes possible. For each of these 2384 acute leukemia patients at least one MLL fusion allele was identified and characterized by sequencing. Several prospective studies were already initiated and first published data verified the reliability of these genomic markers for MRD monitoring.<sup>4</sup> Therefore, the use of these MRD markers will contribute in the future to a better stratification of leukemia patients, which will help to further improve the outcome. In particular, for infant ALL patients, due to the relatively low numbers of potential IG/TR MRD-PCR targets, the availability of an MLL fusion DNA rearrangement has a high impact for the clinical application of MRD monitoring.

The analysis of the MLL recombinome allows to classify MLL fusion partner genes into functional categories. As discussed above, only very few TPGs are recurrently identified with a significant frequency. On the basis of the present study these TPGs are AF4, AF6, AF9, AF10, ELL and ENL. At least for the AF4, AF9, ENL and AF10 proteins exist a functional correlation, as all these proteins are organized within a protein complex (or different subcomplexes) that affect transcriptional elongation. AF4 is the docking platform for AF9 or ENL, which both interact (via AF10) to DOT1L.55,56 DOT1L enable methylation of lysine-79 residues of histone H3 proteins, a prerequisite for the maintenance of RNA transcription<sup>57,58</sup> AF4 binds with its N-terminal portion to the P-TEFb kinase that phosphorylates the largest subunit of RNA polymerase II, DSIF, the NELF complex and UBE2A. This converts RNA POL A into POL E and allows gene transcription.<sup>59</sup> As a result, increased and extended H3K79 methylation signatures seem to accompany the presence of several fusion proteins (MLL-AF4, AF4-MLL, MLL-AF9, MLL-ENL, MLL-AF10 and MLL-AF6),<sup>60</sup> while an additional increase in H3K4 methylation was only demonstrated by the presence of the reciprocal AF4-MLL<sup>59</sup> that causes proB ALL in C57Bl6 mice<sup>61</sup> and was shown to cooperate with the RUNX1 protein.<sup>62</sup> Thus, all the major MLL fusions share a common pathway, which is not only functionally related but offers new and interesting venues to develop new drugs against these leukemias, for example, by the development of DOT1L inhibitors.<sup>63</sup> The fusion proteins MLL-ENL, MLL-AF9 and MLL-AF10 recruit thereby the AF4 complex, while the reciprocal AF4–MLL fusion protein is able to perform exactly the same actions on RNA polymerase II and DOT1L. Thus, future therapies addressing either the inhibition of DOT1L, P-TEFb, or blocking the interaction of the MLL N terminus with MENIN1/LEDGF/MYB are promising new ways to address these leukemias. In addition, the inhibition of Taspase1 would help to inactivate the AF4-MLL fusion protein, as the uncleaved fusion protein is rapidly degraded by SIAH1 and SIAH2.64

For all the other MLL fusion partners identified so far, a systematic classification about their function(s) has been described

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in great detail recently.<sup>65</sup> However, further functional studies are necessary to study the mechanisms that are involved or causative for their leukemogenic activity. Such studies will provide the basis for developing new therapeutic strategies in the future.

# **CONFLICT OF INTEREST**

The authors declare no conflict of interest.

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